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Bad03254 Oryza sat
Bad03677 oryza sat
Q96686 homo sapien
Q95668 homo sapien
Q8556 mycobacteri
Q870y5 neurospora
Q7xp55 oryza sativ
Q9430 streptomyce
Q961b2 pseudomonas
Q9813 rhizobium 1
Q9qud1 tt virus. o
Q9qud1 tt virus. o
Q9qud2 tt virus. o
Q9qud3 xanthomonas
Q8239 streptomyce
Q7pct4 anopheles
Q8pcg3 xanthomonas
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Q726b3 desulfovibr
Aa897722 desulfovi
Q78ad5 neurospora
Q8rnz2 pseudomonas
Q6h4i8 oryza sativ
Q72kt8 thermus the
Aa880777 thermus the
Q6zf42 oryza sativ
Bac83391 oryza sati
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                                                                                                       January 31, 2005, 18:02:30; Search time 105.833 Seconds (without alignments) 54.366 Million cell updates/sec
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GenCore version 5.1.6
Copyright (c) 1993 - 2005 Compugen Ltd.
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0736B3

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Listing first 45 summaries
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Q87x57 pseudomonas Q81x19 phocoena ap Q61ck5 homo sapien Cag30249 homo sapien Q87kt8 tt virus. h Q872kt8 tt virus. o Q94t85 tt virus. o Q94t86 tt virus. o Q94t87 tt virus. o Q94t87 tt virus. o Q94t91 tt virus. o Q94t91 tt virus. o		th / ATCC 29579 / NCIMB
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68.0 68.0 68.0 68.0 68.0 68.0 68.0 68.0		o vulgaris (s oteobacteria; onaceae; Desu 182; M N.A.
32 334 334 334 335 337 337 40 40 117 42 117 44 43 117 45 117	SULT 1 SPIG O98PIG D1-OCT-2001 (TrEMBLrel. 01-OCT-2001 (TrEMBLrel. 01-JUN-2003 (TrEMBLrel. Hypothetical protein MYF OrderedLocusNames=MYPU Mycoplasma pulmonis. Bacteria; Firmicutes; MC NCBL TaxID=2107; [1] SEQUENCE FROM N.A. STRAIN=UAB CTIP; MEDLINE=21267165; PubMec Chamband I., Heilig R., Moszer I., Dybvig K., WI Blanchard A.; The complete genome seq Mycoplasma pulmonis."; Nucleic Acids Res. 29:21 BMBL; H90603; H90603. Complete proteome; Hypot SEQUENCE 72 AA; 7389 Query Match Best Local Similarity 28. Matches 2; Conservative Mycoplasma PRELIMINARY; O72683 O72683 O72683 O72683 O72681 O5-JUL-2004 (TrEMBLrel. O5-JUL-2004 (TrEMBLrel.	Desulfovibrio vulgas 8303). 8303). Bacteria; Proteobaci Desulfovibrionaceae. NCBI_TaxID=882; [1] SEQUENCE FROM N.A. PubMed=15077118; DO
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NCBI_TaxID=5141;
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Heidelberg J.F., Seshadri R., Haveman S.A., Hemme C.L., Paulsen I.T., Kolonay J.F., Eisen J.A., Ward N.L., Mathe B.A., Brinkac L.M., Daugherty S.C., DeBoy R.T., Dodson R.J., Durkin A.S., Madupu R., Nelson W.C., Sullivan S.A., Fouts D.E., Haft D.H., Selengut T., Peterson J.D., Davidsen T.M., Zafar N., Zhou L., Radune D., Davidsen T.M., Zafar N., Zhou L., Radune D., Tran K., Khouri H.M., Gill J., Utterback T.R., "The genome sequence of the anaerobic, sulfate-reducing bacterium Desulfovibrio vulgaris Hildenborough.";

MAC. Biotechnol. 22:554-559(2004).

TIGR; DVU3322; ASS97722.1;
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Mat. Blotechnol. 22.554-559(2004).

EMBL ARONJECT D.
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DVU3252.
Desulfovibrio vulgaris (strain Hildenborough / ATCC 29579 / NCIMB
                                                                                                                                                                                                                                                                                                                                                                                                              Bacteria; Proteobacteria; Deltaproteobacteria; Desulfovibrionales;
Desulfovibrionaceae; Desulfovibrio.
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Pred. No. 6.6e+03;
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Pred. No. 6.6e+03;
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83 AA; 9109 MW;
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Best Local Similarity 28.6%;
Matches 2; Conservative
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The 2, Conservative
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Galagan J.E., Calvo S.E., Borkovich K.A., Selker E.U., Read N.D., Jaffe D., FitzHugh W., Ma L.-J., Smirnov S., Purcell S., Rehman B., Biting T., Engels R., Wang S., Nielsen C.B., Butler J., Endrizzi M., Alting T., Engels R., Wang S., Nielsen C.B., Butler J., Endrizzi M., Cui D., Ianakiev P., Pedersen D., Nelson M., Mashburne M., Selitrennikoff C.P., Kinsey J.A., Braun E.L., Zelter A., Schulte U., Rothe G.O., Jedd G., Mewes W., Staben C., Marcotte E., Greenberg D., Roy A., Foley K., Naylor J., Thomann N., Barrett R., Gnerre S., Kamal M., Kamvysselis M., Maucell E., Bielke C., Rudd S., Frishman D., Krystofova S., Rasmussen C., Metzenberg R.L., Perkins D.D., Kroken S., Cogoni C., Macino G., Catcheside D., Li W., Pratt R.J., Osmani S.A., DeSouza C.C., Glass L., Orbach M.J., Berglund J., Woelker R., Aramayo R., Yarden O., Plamann M., Seiler S., Dunlap J., Radford A., Aramayo R., Paulsen I., Sachs M.S., Lander E.S., Nusbaum C., Birren B.; The Genome Sequence of the Filamentous Fungus Neurospora crassa."; Nature 0:0-0 (2003).
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"A Functional Screen for the Type III (Hrp) Secretome of the Plant
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     -1- CAUTION: The sequence shown here is derived from an EMBL/GenBank/DDBJ whole genome shotgun (WGS) entry which is
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01-JUN-2002 (TrEMBLrel. 21, Last sequence update)
01-JUN-2002 (TrEMBLrel. 21, Last annotation update)
Putative type III effector HolPtov (Fragment).
Pseudomonas syringae (pv. tomato).
Bacteria; Proteobacteria; Gammaproteobacteria; Pseudomonadales;
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Neurospora crassa.
Eukaryota; Fungi; Ascomycota; Pezizomycotina; Sordariomycetes;
Sordariomycetidae; Sordariales; Sordariaceae; Neurospora.
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VCE 87 AA; 8712 MW; 19B486AC40439088 CRC64;
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EMBL; AF458401; AAL84265.1; -...
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TTCU429.
Thermus thermophilus (strain HB27 / ATCC BAA-163 / DSM 7039).
Bacteria, Deinococcus-Thermus, Deinococci, Thermales, Thermaceae;
  14-APR-2004 (TrEMBLrel. 27, Created)
14-APR-2004 (TrEMBLrel. 27, Last sequence update)
11-MAY-2004 (TrEMBLrel. 27, Last annotation update)
Hypothetical membrane spanning protein.
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                                                                                                                                                                                          Oryza sativa (japonica cultivar-group).
Bukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae;
Ehrhartoideae; Oryzeae; Oryza.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Thermus thermophilus (strain HB27 / ATCC BAA-163 / DSM 7039).
Bacteria, Deinococcus-Thermus, Deinococci, Thermales, Thermaceae,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Henne A., Brueggemann H., Raasch C., Wiezer A., Hartsch T., Liesegang H., Johann A., Lienard T., Gohl O., Martinez-Arias R., Jacobi C., Starkuviene V., Schlenczeck S., Dencker S., Huber R., Klenk H.-P., Kramer W., Merkl R., Gottschalk G., Fritz H.-J.; "The genome sequence of the extreme thermophile Thermus thermophilus.";
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           5; Indels
                                                                                                                                                                                                                                                                                                                                              Sasaki T., Matsumoto T., Katayose Y.,
Submitted (OCT-2002) to the EMBL/GenBank/DDBJ databases.
EMBL; AP005867; BAD26361.1; -
                                                                                                                                                                                                                                                                                                                                                                                                           Hypothetical protein.
SEOUBNCE 102 AA; 10257 MW; 2330B2E1FB71DDEC CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Complete proteome; Hypothetical protein.
SEQUENCE 109 AA; 11814 MW; B143CD19CA62593F CRC64;
                                                                                05-JUL-2004 (TrEMBLrel. 27, Created)
05-JUL-2004 (TrEMBLrel. 27, Last sequence update)
05-JUL-2004 (TrEMBLrel. 27, Last annotation update)
Hypothetical protein P0599F09.8.
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05-JUL-2004 (TrEMBLrel. 27, Last sequence update)
05-JUL-2004 (TrEMBLrel. 27, Last annotation update)
Hypothetical membrane spanning protein.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                            68.0%; Score 17; DB 2; I
28.6%; Pred. No. 7.8e+03;
                                               102 AA
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EMBL, AEU17302, AAS80777.1; -.
InterPro; IRR005530; SPW.
Pfam; PF03779; SPW; 2.
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Best Local Similarity 28.6'
....has 2; Conservative
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RESULT 6
264418
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AAS80777
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Eukaryota, Viridiplantae, Streptophyta, Embryophyta, Tracheophyta,
Spermatophyta, Magnoliophyta, Liliopsida, Poales, Poaceae,
Henne A., Brueggemann H., Raasch C., Wiezer A., Hartsch T.,
Liesegang H., Johann A., Lienard T., Gohl O., Martinez-Arias R.,
Jacobi C., Starkuviene V., Schlenczeck S., Dencker S., Huber R.,
Klenk H.-P., Kramer W., Merkl R., Gottschalk G., Fritz H.-J.;
"The genome sequence of the extreme thermophile Thermus
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28.6%; Pred. No. 8.3e+03;
ative 0; Mismatches 5; Indels
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28.6%; Pred. No. 9.4e+03;
Live 0; Mismatches 5; Indels
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Submitted (OCT-2001) to the EMBL/GenBank/DDBJ databases.
EMBL; AP004270; BAC83391.1; -.
Hypothetical protein.
SEQUENCE 127 As; 13838 MW; 6F74D278AC08AF0C CRC64;
                                                                                                                                               Nat. Biotechnol. 22:547-553(2004).
EMBL; AE017302; AAS80777.1; -.
Hypothetical protein.
SEQUENCE 109 AA; 11814 MW; B143CD19CA62593F CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            05-JUL-2004 (TrEMBLrel. 27, Created)
05-JUL-2004 (TrEMBLrel. 27, Last sequence update)
05-JUL-2004 (TrEMBLrel. 27, Last annotation update)
Hypotheital protein P0406F06.31.
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02-MAR-2004 (TrEMBLrel. 27, Last sequence update)
02-MAR-2004 (TrEMBLrel. 27, Last annotation update)
Hypothetical protein P0406F06.31.
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Matches 2; Conservative
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ID BAC8
AC BAC8
DT 02-N
DT 02-N
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This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation - the European Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/or send an email to license@isb-sib.ch).
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Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae;
Ehrhartoideae; Oryzeae; Oryza; Oryza sativa.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Urabe H., Ogawara H.; "Nucleotide sequence and transcriptional analysis of activator-regulator proceins for beta-lactamase in Streptomyces cacaoi."; J. Bacteriol. 174:2814-2842(1992).
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                                                                        SEQUENCE FROM N.A.
SEQUENCE FROM N.A.
STRAIN=cv. Nipponbare;
Sasaki T., Macaumoto T., Yamamoto K.;
Sasaki T., Macaumoto T., Yamamoto K.;
Supari Sativa nipponbare (GA3) genomic DNA, chromosome 7, PAC clone:P0406F06.";
Submitted (OCT-2001) to the EMBL/GenBank/DDBJ databases.
EMBL; AP004270; BAC83391.1; -
Hypothetical protein.
SEQUENCE 127 AA; 13838 MW; 6F74D278AC08AF0C CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Bacteria; Actinobacteria; Actinobacteridae; Actinomycetales;
Streptomycineae; Streptomycetaceae; Streptomyces.
NCBI_TaxID=1898;
                                                                                                                                                                                                                                                   Length 127;
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Pred. No. 9.4e+03;
0; Mismatches 5; Indels
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28.6%; Pred. No. 9.7e+03;
tive 0; Mismatches 5; Indels
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PIR; C41855; C41855.
Hypothetical protein.
SEQUENCE 132 Aa; 14234 MW; FFD484A2182B52D8 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         (TrEMBLrel. 27, Created)
(TrEMBLrel. 27, Last sequence update)
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01-FEB-1994 (Rel. 28, Last sequence update)
01-FEB-1994 (Rel. 28, Last annotation update)
Hypothetical 14.2 kDa protein in blaB 3 region.
                                                                                                                                                                                                                                                                                                                                                                                                                                 132 AA.
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MEDLINE=92234939; PubMed=1569015;
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28.6%;
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Best Local Similarity 28.00,
2; Conservative
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                                                   NCBI_TaxID=39947;
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05-JUL-2004
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Q6Z0D1
ID Q6Z0D1
AC Q6Z0D1
DT 05-JUI
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Gaps
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                                                P0703C03.23).
Name-CSJNBa0078D03.43; Synonyms=P0703C03.23;
Name-CSJNBa0078D03.43; Synonyms=P0703C03.23;
Sylvidiplanca cultivar-group).
Bukaryota; Viridiplancae; Streptophyta; Embryophyta; Tracheophyta; Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae; Ehrhartoideae; Oryzeae; Oryzea.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               02-MAR-2004 (TrEMBLrel. 27, Created)
02-MAR-2004 (TrEMBLrel. 27, Last sequence update)
02-MAR-2004 (TrEMBLrel. 27, Last annotation update)
02-MAR-2004 (TrEMBLrel. 27, Last annotation update)
14-Mypothetical protein Po703C03.23.

Oryza sativa (japonica cultivar-group).

Bukaryota, Viridiplantae, Streptophyta; Embryophyta; Tracheophyta; Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae;
Bhrhartoideae; Oryzeae; Oryza; Oryza sativa.
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05-JUL-2004 (TrEMBLrel. 27, Last annotation update)
Hypothetical protein OSJNBa0078D03.43 (Hypothetical protein
P0703C03.23).
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Clone:P0703C03.";
Submitted (JAN-2002) to the EMBL/GenBank/DDBJ databases.
EMBL; AP004G37; BAD03254.1; -.
Hypothetical protein.
SEQUENCE 133 AA; 14574 MW; 9F18BC7F0A45B2BA CRC64;
                                                                                                                                                                                                                                                                  Sasaki T., Matsumoto T., Yamamoto K.;
Submitted (JAN-2002) to the EMBL/GenBank/DDBJ databases.
EMBL; AP005493; BAD03677.1; -.
EMBL; AP006437; BAD03254.1; -.
Hypothetical protein.
SEQUENCE 133 AA; 14574 MW; 9F18BC7F0A45B2BA CRC64;
                                                                                                                                                                                 SEQUENCE FROM N.A.
Sasaki T., Matsumoto T., Katayose Y.;
Submitted (JUL-2002) to the EMBL/GenBank/DDBJ databases
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02-MAR-2004 (TEMBLrel. 27, Created)
02-MAR-2004 (TEMBLrel. 27, Last sequence update)
02-MAR-2004 (TEMBLrel. 27, Last annotation update)
Hypothetical protein OSJNBa0078D03.43.
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Sasaki T., Matsumoto T., Yamamoto K.;
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Best Local Similarity 28.00,
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ses 2; Conservative
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BAD03254;
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Klausner R.D., Collins F.S., Wagner L., Shenmen C.M., Schuler G.D.,
Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
Boatchenko L., Marusina K., Farmer A.A., Rubin G.M., Hong L.,
Brapleton M., Soarse M.B., Bonaldo M.P., Casavant T.L., Scheetz T.E.,
Brownstein M.J., Usdin T.B., Toshiyuki S., Carninci P., Frange C.,
Raha S.S., Loquellano N.A., Peters G.J., Abramson R.D., Mullahy S.J.,
Brichards S., Worley K.C., Hale S., Garcia A.M., Gay L.J., Hulyk S.W.,
Villalon D.K., Muzny D.M., Sodergren E.J., Lu X., Gibbs R.A.,
Whiting M., Madan A., Young A.C., Shevychenko Y., Bouffard G.G.,
Blakesley R.W., Touchman J.W., Green E.D., Dickson M.C.,
Blakesley R.W., Touchman J.W., Green E.D., Dickson M.C.,
Jones S.J., Marra M.A.,
Jones G.J., Marra M.J.,
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             Oryza sativa (japonica cultivar-group).
Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;
Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae;
Ehrhartoideae; Oryzeae; Oryza; Oryza sativa.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi,
Mammalia, Eutheria, Primates, Catarrhini, Hominidae, Homo.
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                                                                                                                                                                                                STRAIN=cv. Nippondare; Sasaki T., Matsupose Y.; Sasaki T., Matsumoto T., Katayose Y.; Sasaki T., Matsumoto T., Katayose Y.; Matsumosasulva nippondare(GA3) genomic DNA, chromosome 8, BAC clone:OSJNBa10078B03."; Submitted (JUL-2002) to the EMBL/GenBank/DDBJ databases. BMBL; AP005493; BAD03677.1; -. Hypothetical procedin. SEQUENCE 133 AA; 14574 MW; 9F18BC7F0A45B2BA CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                  Score 17; DB 2; Length 133;
Pred. No. 9.8e+03;
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Straubberg R.;
Submitted (AUG-2001) to the EMBL/GenBank/DDBJ databases.
EMBL; BC012317; AAH12317.1; -.
Hypothetical protein.
SEQUENCE 137 AA; 14308 MW; 53A2EF0CB2BCFB78 CRC64;
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01-DEC-2001 (TrEMBLrel. 19, Last sequence update)
01-MAR-2004 (TrEMBLrel. 26, Last annotation update)
Hypothetical protein LOC93082.
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MEDLINE-22388257; PubMed-12477932;
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28.6%;
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Length 137;

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Query Match

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             5; Indels
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